REARING OF *DIATRAEA SACCHARALIS* ON DIETS IN SURINAM

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BY

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Mass rearing of the borer *Diatraea saccharalis* (F.) was attempted on a limited scale. Out of nine non-aseptic artificial diets tested fair results were obtained with the medium containing kidney bean meal (35 g), corn plant powder (10 g), carrot powder (5 g), brewer's yeast (10 g), casein (1 g), ascorbic acid (0.5 g), agar (3 g), water (210 g) and the antimicrobial agents methylparahydroxybenzoate (1 g), streptomycine sulphate (0.1 g), penicillin (0.2 g) and sorbic acid (0.2 g).

The borer was also reared satisfactorily on cut corn stalks.

*Diatraea saccharalis* is one of the most important insect pests attacking sugarcane and rice in Surinam. Control by insecticides has not proved economical or feasible because the larvae live predominantly inside the plant stalks and rainfall limits or eliminates the residual effect of the insecticides (Van Dinther, 1960). As a result of this and also in view of the general objections against the application of chemicals, interest is now centred on the possibilities of biological control.

Good results have been obtained with the introduction of parasites of this borer in a number of areas of the Caribbean since the pioneer work of Myers (1934, 1935) and Box (1939 a, b). For example, on the 8000 hectare sugarcane estate "El Palmar" in the Aragua Valley, Venezuela, after the liberation of the "Amazon fly", *Metagonistylum minense* Tns., the mean stalk infestation by *D. saccharalis*, *D. rosa* Heinr. and *D. busckella* Dyar & Heinr. gradually decreased from 16% in 1947—1950 to 2% in 1968. At this estate the fly is mass-reared and about 60,000 are released annually during May-November (Dr. José Morejon, head of the breeding and release work, pers. comm.).

By contrast, in several other countries parasite establishment was not successful. For a general review of the biological control status of moth borers reference can be made to Jepson (1954), Simmonds & Bennett (1967) and Bennett (1969).

Envisaging the eventual introduction into Surinam of exotic parasites, preliminary research on an economically practicable mass-rearing method of the borer was started in 1968 and attention was especially directed to devising artificial diets.

Different artificial diets for lepidopterous larvae have developed in the U.S.A.

In Surinam we tested a number of media, based on the simple recipe of Shorey & Hale. Some new compositions and ingredients were examined. The possibility of mass rearing *D. saccharalis* on more natural food, viz. on cut rice and corn stalks, was also studied.

**DIETS AND PREPARATION**

Details of the components of the diets, listed in Table I, are as follows: Initially dried kidney beans (*Phaseolus* sp.), were soaked in water overnight, then boiled and ground into pulp with a handmincer. Later the beans were applied in the form of a fine powder, made by the Department of Food Technology, Agricultural Experiment Station, Paramaribo, who also provided corn-, rice- and carrot-powder. The orginal plant materials — viz. stalks and leaves of corn- and rice-plants that had not yet reached the generative stage and imported deep-frozen roots of *Daucus carota* — were slowly dried at 45° C before grinding to preserve the vitamins and aromatics in them.

Brewer’s yeast, casein, (cane) sugar, ascorbic acid, choline chloride and agar powder are among the ingredients already included in the media mentioned earlier. To prevent contamination two or more of the following antibiotics (“inhibitors”)

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**TABLE I**

<table>
<thead>
<tr>
<th>Diet No.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
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<th>6</th>
<th>7</th>
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<td>15</td>
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<tr>
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<td>5</td>
<td>15</td>
<td>5</td>
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</tr>
<tr>
<td>rice plant powder</td>
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<td>10</td>
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<td>6</td>
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<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
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<td>0.5</td>
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<tr>
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<tr>
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<tr>
<td>copper sulphate</td>
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<tr>
<td>Dimanin</td>
<td>0.5</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>water</td>
<td>375</td>
<td>375</td>
<td>190</td>
<td>210</td>
<td>190</td>
<td>280</td>
<td>210</td>
<td>200</td>
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</table>
generally applied in artificial diets, were used: methylparahydroxybenzoate (nipagin), streptomycin, sorbic acid and formaldehyde. Penicillin, copper sulphate, Dimanin (trade-mark for a mixture of metal salts with fungicidal and bactericidal action) and hydrochloric acid (1 N) were also tested.

Diet 1 was prepared as follows: brewer's yeast, ascorbic acid, nipagin and streptomycin in 250 cc of water were blended in a beaker for 15 sec with an electric mixer. The agar, dissolved in a remaining 125 cc of water at 40°, was poured over this mixture and the freshly boiled and ground beans, the corn meal and the carrot powder were added. All ingredients together were stirred for 30 sec. The medium was then poured into a plastic squeeze bottle and dispensed into rearing cups and vials. These containers were covered with paper and left open for one day to let the excess moisture evaporate. In this medium, sorbic acid and formaldehyde were omitted but streptomycin was added as a precaution to prevent rotting. The diet was enriched with corn meal and carrot powder as recommended by Walker et al., (1966).

In composing diet 2 special attention was paid to the keeping qualities. Copper sulphate, Dimanin and hydrochloric acid were added as extra bacterial and fungal inhibitors. The agar was dissolved in water at 90°, and brewer's yeast, corn meal and carrot powder added while stirring. At 70° the remaining constituents followed and the medium was mixed for 2 min. The cups and vials were briefly submerged in alcohol before use. After the hot medium was poured into these containers they were covered overnight with paper disinfected in alcohol.

Diet 3 comprised the ingredients used by Shorey & Hale (1965) and also some of the components of the wheat germ medium of Adkisson et al., (1960). Hydrochloric acid, copper sulphate and Dimanin were abandoned; streptomycin and penicillin supplemented the other antimicrobial agents.

In the diets tested subsequently bean meal was used. The handmixer was replaced by a blender with a 1 l beaker. The dry ingredients, except the agar, were mixed with about half of the available volume of water for 30 sec. The agar was dissolved in the remaining water at 90°. When cooled to 70° it was added to the mixture which was blended for another 1—2 min.

REARING PROCEDURE

The breeding work was carried out in a screened laboratory room where the mean temperature fluctuated from 24°—28° and the relative humidity ranged from about 65—90%. Daylength was approximately 12 hours and no extra light was supplied.

*D. saccharalis* pupae, collected from rice fields at Paramaribo, formed the initial material. They were placed on a layer of moistened cotton in an open petri-dish and kept in a cardboard cylinder (height 30 cm, diameter 23 cm) the inner wall of which was lined with ordinary writing-paper. The open top was covered by the same type of paper whereas a 29 cm high and zigzag folded sheet of paper was put inside. A small wad of cotton soaked in a sugar solution was also supplied.
Moths mated soon after emergence and egg-clusters were readily deposited on the writing-paper. Before hatching the egg-groups were cut out from the paper sheets. When not directly needed, eggs were kept in a refrigerator (6°). If freshly deposited they can be preserved in a good condition for a few days only; eggs in the "blackhead stage" can be stored up to 10 days. Pupae were kept in a similar way.

Newly hatched larvae were transferred by means of a fine brush (no. 00) to cotton wool plugged glass vials (length 10 cm, diameter 2.5 cm) containing a medium. The brush was regularly dipped into a 0.1% solution of the antisepticum "Dettol" (chloroxylenol). Initially 25 cc plastic cups with lid were used. To assure a sufficient air supply a 2 cm wide hole was punched in the lid and stoppered with cotton wool. However, the use of these cups was soon abandoned because larvae may gnaw an opening in the wall.

In the experiments with stalks of rice and corn 20 cm long sections, cut from plants that had not reached the flowering stage, were kept in 2 1 glass jars. The 8 cm wide jar mouth was closed by a pierced wooden stopper, covered with a fine mesh gauze. One or two egg-masses were deposited on this plant material.

To ascertain the fecundity of individual females, couples of freshly emerged moths were kept in paper cups (height 20 cm, diameter 10 cm) where eggs were laid and counted.

RESULTS AND DISCUSSION

The biological data obtained have been recorded in Table II. Some diets, successfully applied elsewhere, failed to yield good results under our conditions. First-instar larvae were mobile during the first days after introduction. In the cups they finally settled in the narrow space along the wall, caused by the shrinking of the medium.

The keeping qualities of diet 1 proved to be poor. One week after larval introduction this medium had changed from light brown to a darker brown and a putrid smell was emitted. Containers revealing a slight infection only were kept for further observation. Notwithstanding the fact that the contamination soon intensified 36% of the larvae reached the pupal stage. The period necessary for the total larval development was strikingly short.

The extra inhibitors used in diet 2 prevented any decomposition but larval development was so much retarded that only a limited number of first-instar larvae had moulted after 9 days. After 6 weeks the observations were concluded when only a few larvae had attained the fourth stage.

The results obtained with diet 3 were much improved after leaving out some of the inhibitors, including Dimanin, but still about half the larvae died.

Good results were produced by medium 4, viz. a rather low larval mortality, a reasonable developmental duration, normal pupal weights and a fair production of viable eggs. Four consecutive generations of D. saccharalis had been reared successfully on this diet when the experiment was topped.
**TABLE II**  
*Biological data of D. saccharalis reared on artificial diets, rice stalks and corn stalks*

<table>
<thead>
<tr>
<th>Diet no.</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>rice stalks</th>
<th>corn stalks</th>
</tr>
</thead>
<tbody>
<tr>
<td>number of larvae tested</td>
<td>150</td>
<td>160</td>
<td>180</td>
<td>50</td>
<td>70</td>
<td>50</td>
<td>70</td>
<td>70</td>
<td>50</td>
<td>100</td>
<td>200</td>
</tr>
<tr>
<td>number of larvae per cup/vial</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td>5</td>
<td>5</td>
<td>6</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>diet weight (g) per cup/vial</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>14</td>
<td>16</td>
<td>16</td>
<td>16</td>
<td>14</td>
<td>14</td>
<td></td>
<td></td>
</tr>
<tr>
<td>larval stage (days) *</td>
<td>23 ± 0.9</td>
<td>&gt;43</td>
<td>38 ± 0.4</td>
<td>32 ± 0.6</td>
<td>&gt;64</td>
<td>41 ± 0.6</td>
<td>40 ± 0.8</td>
<td>43 ± 1.2</td>
<td>44 ± 1.1</td>
<td>30 ± 0.9</td>
<td>28 ± 0.7</td>
</tr>
<tr>
<td>larval mortality (%)</td>
<td>64</td>
<td>40</td>
<td>48</td>
<td>19</td>
<td>75</td>
<td>27</td>
<td>44</td>
<td>61</td>
<td>36</td>
<td>65</td>
<td>30</td>
</tr>
<tr>
<td>avg. pupal stage (days)</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>7</td>
<td>7</td>
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</tr>
<tr>
<td>pupal weight (mg) *</td>
<td>49 ± 2.5</td>
<td>57 ± 0.9</td>
<td>59 ± 1.1</td>
<td>42 ± 1.1</td>
<td>39 ± 1.5</td>
<td>45 ± 2.1</td>
<td>47 ± 2.5</td>
<td>55 ± 1.5</td>
<td>46 ± 0.7</td>
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<tr>
<td>of males</td>
<td>68 ± 4.0</td>
<td>93 ± 1.1</td>
<td>112 ± 3.1</td>
<td>89 ± 2.3</td>
<td>91 ± 2.3</td>
<td>87 ± 4.2</td>
<td>94 ± 3.3</td>
<td>99 ± 2.9</td>
<td>103 ± 2.0</td>
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</tr>
<tr>
<td>egg-production/♀ *</td>
<td>308 ± 11</td>
<td>394 ± 2.5</td>
<td>316 ± 13</td>
<td>274 ± 15</td>
<td>267 ± 16</td>
<td>253 ± 24</td>
<td>418 ± 35</td>
<td>453 ± 27</td>
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<tr>
<td>avg. egg stage (days)</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>7</td>
<td>7</td>
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<td>7</td>
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<tr>
<td>egg-viability (%)</td>
<td>92</td>
<td>94</td>
<td>85</td>
<td>80</td>
<td>79</td>
<td>69</td>
<td>99</td>
<td>97</td>
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</table>

* * mean and standard deviation of the mean
The effect on borer growth by diet 5, a copy of Bowling’s (1967) recipe with only the “pinto beans” replaced by kidney beans, was disappointing. Larval development was extremely slow and mortality high among the first three instars. When the observations were terminated after 9 weeks only 25% of the larvae were still alive. This result was probably due to the pasty texture of the medium.

The outcome of diet 6, mainly composed of constituents applied by Walker et al., proved to be moderate.

Diet 7 showed some resemblance to the medium of Pan & Long (1961). Instead of autoclaving to sterilize the medium extra fungicides were used. Again the speed of larval growth was slow and mortality high. The food consistency appeared to be too compact and limited the boring activities of the larvae.

The last two experiments conducted with artificial diets demonstrated that rice-plant powder and rice polishings may replace to some extent corn meal and beans or corn meal and carrot-powder. However, the tardy growth and the high larval mortality render these diets impracticable. Thus medium 4 was the most promising diet offered.

Mass-breeding on cut rice stalks was unsatisfactory because they needed replacing every 4 to 5 days. The use of corn stalks was less laborious as they had to be renewed only once every 14 days. Since corn can be grown throughout the year and reasonable rearing results were obtained (see Table II) the breeding of D. saccharalis on corn needs further evaluation before a final choice of artificial diet is made.

Résumé

ÉLEVAGE DE DIATRAEA SACCHARALIS SUR MILIEUX ARTIFICIELS EN SURINAM

L’élevage de Diatraea saccharalis sur une huitaine de régimes artificiels a été étudié sous des conditions non-aseptiques. Des résultats favorables sont obtenus avec un milieu composé de: haricots rouges pulvérisées (35 g), tiges et feuilles de maïs réduites en poudre (10 g), farine de carottes (5 g), levure (10 g), caséine (1 g), acide ascorbique (0,5 g), agar (3 g), eau (210 g) et les antibiotiques: méthylparahydroxybenzoate (1 g), streptomycine sulfate (0,1 g), pénicilline (0,2 g) et acide sorbique (0,2 g).

L’élevage en bocaux contenant des fragments de tiges de maïs a également donné de bons résultats.

Références


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